

New Electrocatalysts For Fuel Cells

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Objective: Reduction of precious metal loading

•DOE Technical Barriers for Fuel Cell Components

- Q. Electrode Performance

•Budget: FY2003: \$ 400 K

FY2004: \$ 450 K

Staff Scientist: Nenad M. Markovic

Post Doctoral Fellow: Vojislav Stamenkovic

Graduate Students: Berislav Blizanac (Belgrade)

Karl Mayrhofer (Vienna Tech. Univ.)

LBNL Materials-by-Design Approach rrrrrBERKELEY LAB Model Systems Real Catalysts Ex-Situ **TEM XPS** LEIS LEED **AES** In-Situ **RRDE FTIR** $4.4 \pm 1.6 \text{ nm}$ 30 **SXS Kinetics** [%] 25 20 15 10 (c) single crystal electrode 5 10



Collaborations

Industries

Characterization and testing of new catalysts for developers

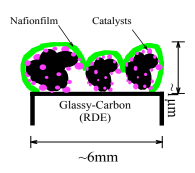
- **►** GM , Rochester, NY, USA
- FIFC, South Windsor, CT, USA
- > 3M, Minneapolis, MN, USA

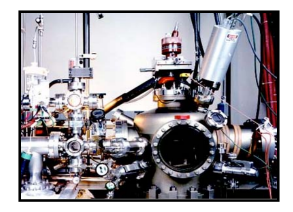
Universities and Institutes

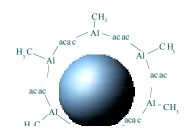
Synthesis of metallic nanoclusters

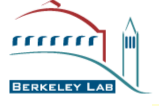
- Max-Planck-Institut fuer Kohlenforschung, Muelheim/Ruhr, Germany
- University of Liverpool, UK











Research Plan: 2003-2004

New catalysts for both anodes and cathodes being developed following a unified concept of PGM-based bimetallic nanoparticles with a "grape" structure (a PGM "skin" with base metal core)

Choice of PGM and core metals different for anode and cathode PGM/base metal combinations based on computational screening of PGM core-shell nanostructures using newly developed (under BES funding) Monte Carlo simulations

Fundamental studies of the crystallite size effect for the oxygen reduction reaction in acidic electrolytes on carbon supported Pt and Pt alloy nanoparticles

Pursue new synthetic chemistry to synthesize nanoparticles with a "grape" structure

Continue focus on Re as metal core with Pt and Pd as PGM

Optimization of AuPd as alternative to Pt in anodes

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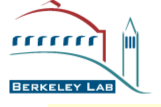
Publications (Since 10/2002)

Refereed Journals and Refereed Conference Proceedings

- 1. Schmidt TJ. Stamenkovic V. Markovic NM. Ross PN. "Electrooxidation of H₂, CO and H₂/CO on well-characterized Au(111)-Pd surface alloys." *Electrochimica Acta*. 48, 3823-3828, 2003 Nov 15.
- 2. Arenz M. Stamenkovic V. Schmidt TJ. Wandelt K. Ross PN. Markovic NM. "The electro-oxidation of formic acid on Pt-Pd single crystal bimetallic surfaces." *Physical Chemistry Chemical Physics.* 5, 4242-4251, 2003 Oct 1.
- 3. Stamenkovic V. Schmidt TJ. Ross PN. Markovic NM. "Surface segregation effects in electrocatalysis: kinetics of oxygen reduction reaction on polycrystalline Pt₂Ni alloy surfaces." *Journal of Electroanalytical Chemistry* 554,191-199, 2003 Sep 15.
- 4. Arenz M. Schmidt TJ. Wandelt K. Ross PN. Markovic NM. "The oxygen reduction reaction on thin palladium films supported on a Pt(111) electrode." *Journal of Physical Chemistry B* 107(36), 9813-9819, 2003 Sep 11.
- 5. Arenz M. Stamenkovic V. Ross PN. Markovic NM. "Preferential oxidation of carbon monoxide adsorbed on Pd submonolayer films deposited on Pt(100)." *Electrochemistry Communications* 5(9), 809-813, 2003 Sep.
- 6. Arenz M. Stamenkovic V. Schmidt TJ. Wandelt K. Ross PN. Markovic NM. "The effect of specific chloride adsorption on the electrochemical behavior of ultrathin Pd films deposited on Pt(111) in acid solution." Surface Science 523, 199-209, 2003 Jan 10.
- 7. Schmidt TJ. Stamenkovic V. Ross PN. Markovic NM. "Temperature dependent surface electrochemistry on Pt single crystals in alkaline electrolyte Part 3. The oxygen reduction reaction." *Physical Chemistry Chemical Physics* 5, 400-406, 2003.
- 8. Stamenkovic V. Schmidt TJ. Ross PN. Markovic NM. "Surface composition effects in electrocatalysis: Kinetics of oxygen reduction on well-defined Pt₃Ni and Pt₃Co alloy surfaces." *Journal of Physical Chemistry B* 106, 11970-11979, 2002 Nov 21.

Books and Book Chapters

- 1. Markovic NM. Radmilovic V. Ross PN. "Physical and Electrochemical Characterization of Bimetallic Nanoparticle Electrocatalysts", in *Catalysis and Electrocatalysis at Nanoparticle Surfaces*, Ed. Wieckowski A. Savinova ER. Vayenas CG., Marcel Dekker, New York and Basel, 2003, Chapter 9, pp. 311-342.
- 2. Ross, PN. "Oxygen Reduction Reaction on Single Crystal Electrodes", in *Handbook of Fuel Cells: Fundamentals, Technology and Applications, Volume 2, Electrocatalysis*, Ed. Viestich W. Lamm A. Gasteiger H., John Wiley & Sons Ltd., Chichester, 2003, pp. 465-481.



Accomplishments Outline

- Results of Monte-Carlo Simulations of Microstructure in Pt₃Ni and Pt₃Re Nanoparticles (DOE/BES sponsored research)
- ➤ High precision activity measurements for well-characterized Pt alloys (i.e. what is the most active alloy and what activity enhancement could we expect)

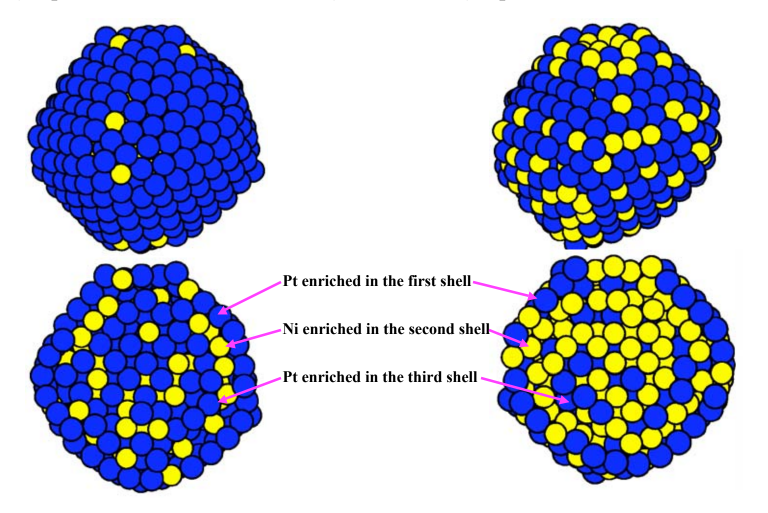
 (GM sponsored research)
- ➤ Definitively determined the leaching out of the transition metal in all Pt₃M alloys in acid electrolyte and its consequences
- Correlated enhanced specific activity of Pt3Co alloy catalyst for ORR to electronic state of Pt skin with Co-enriched second layer
- ➤ Re-examined the "crystallite size effect" for the oxygen reduction reaction on carbon supported Pt nanoparticles
- ➤ Measured surface area and activity of 3M "nanostructured" (NS) catalysts and benchmarked against conventional carbon supported Pt catalyst



Surface-sandwich Structures of Pt-Ni Nanoparticles

Pt₇₅Ni₂₅ fcc cubo-octahedral nanoparticle (snapshot and [001] cross-section)

Pt₅₀Ni₅₀ fcc cubo-octahedral nanoparticle (snapshot and [001] cross-section)



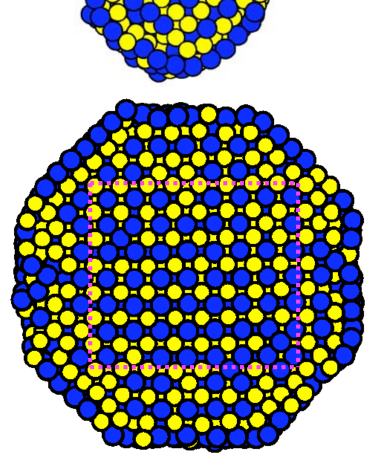


Nanoparticle Structures and Order-Disorder Transitions

Segregation profiles (in atomic concentrations of Pt atoms) of equilibrium cubo-octahedral Pt₅₀Ni₅₀ nanoparticles simulated at T=600K

N	C_1	C_2	C_3	C_{core}
586	70	27	44	35 /
1289	74	31	43	35
2406	79	36	38	37
4033	81	37	41	39

Surface-sandwich structure with a disordered core for smaller nanoparticles



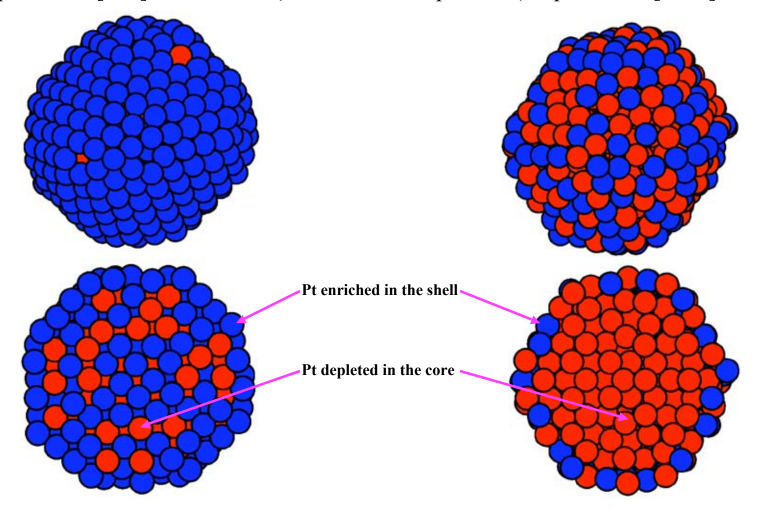
Core-shell structure with an ordered core for larger nanoparticles



Core-shell Structures of Pt-Re Nanoparticles

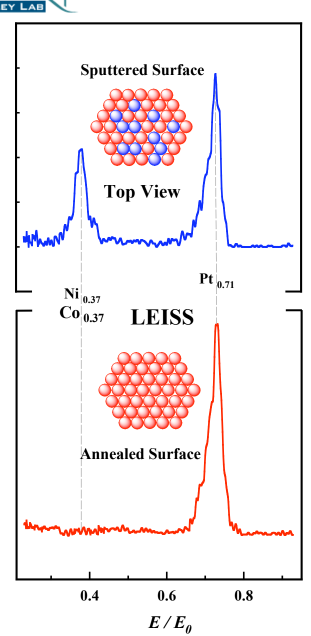
Pt₇₅Re₂₅ fcc cubo-octahedral nanoparticle (snapshot and [001] cross-section)

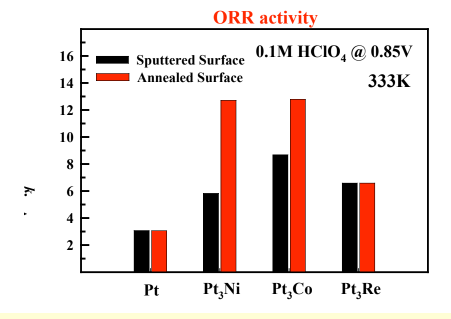
 $Pt_{25}Re_{75}$ hcp truncated hexagonal bipyramidal nanoparticle (snapshot and $[11\overline{2}0]$ cross-section)





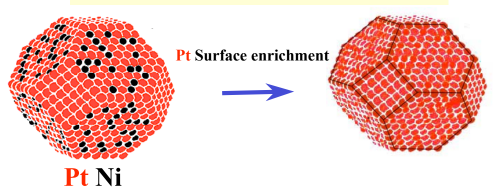
Segregation Effect: Platinum Skin vs. Bulk Alloy Surfaces





Platinum Skin Effect: Bimetallic Nanoparticle

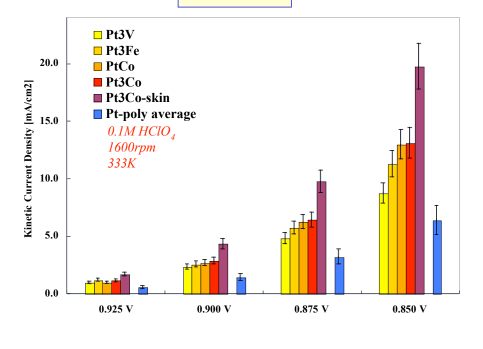
➤ Higher intrinsic activity (per unit area)➤ Substitution of "buried" Pt atoms in particle core by base metal atoms



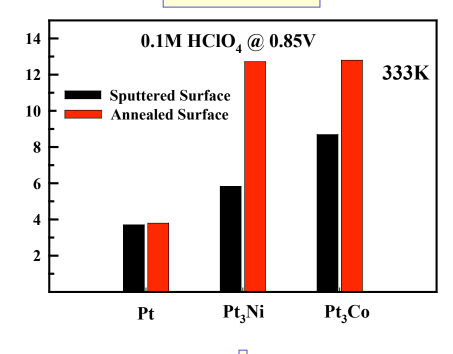
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Activity Factors





Published results



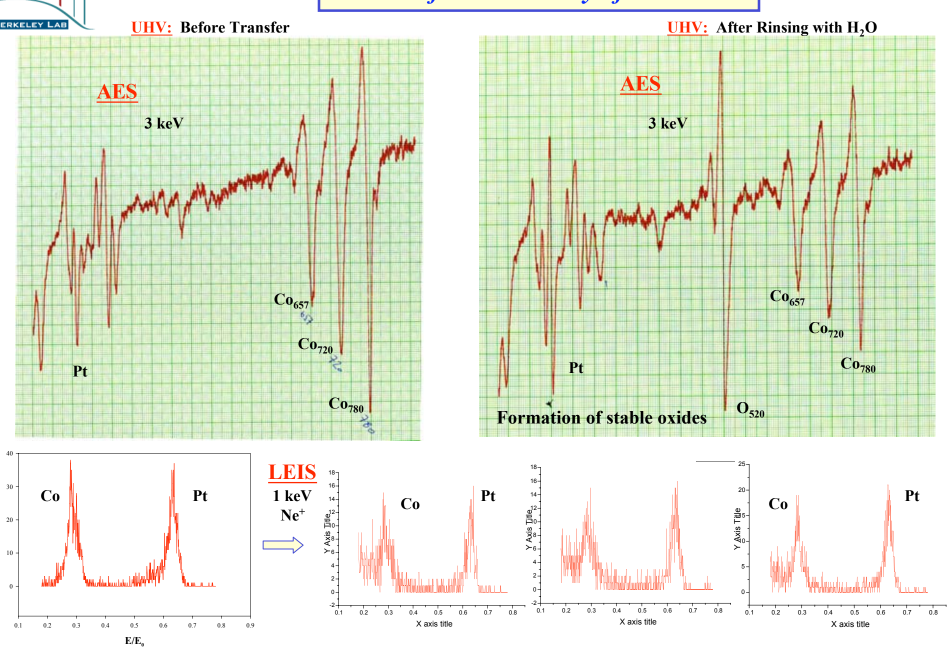
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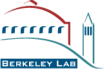
Pt ₃ V	Pt ₃ Fe	PtCo	Pt ₃ Co	Pt ₃ Co skin
1.4 1.0-1.8	1.8 1.4-2.4	2.0 1.5-2.7	2.0 1.5-2.7	3.1 2.3-4.2

Pt poly	Pt ₃ Ni	Pt ₃ Co	Pt ₃ Ni skin	Pt ₃ Co skin
ı	1.6	2.3	3.2	3.4

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Surface Chemistry of PtCo

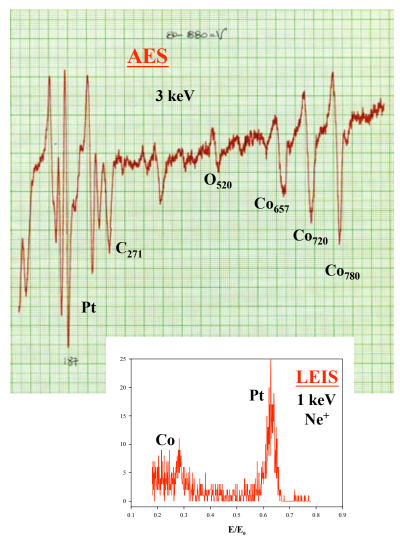


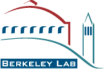


Surface Chemistry of PtCo

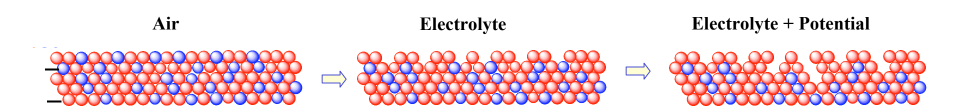
UHV: Before Transfer **AES** 3 keV Pt Co₆₅₇ Co₇₂₀ **LEIS** Co₇₈₀ Pt 1 keV Co Ne^+ 0.3 0.8 0.5 E/E

<u>UHV:</u> After Electrochemistry and/or After rinsing with 0.1M HClO₄





Effect of Dissolution of Alloying Metal



Could active surface area (Pt sites) be higher vs. Pt-poly due to dissolution of alloying components?

$\mathbf{H}_{\mathbf{u}\mathbf{p}\mathbf{d}}$ charge as measure of Pt site density

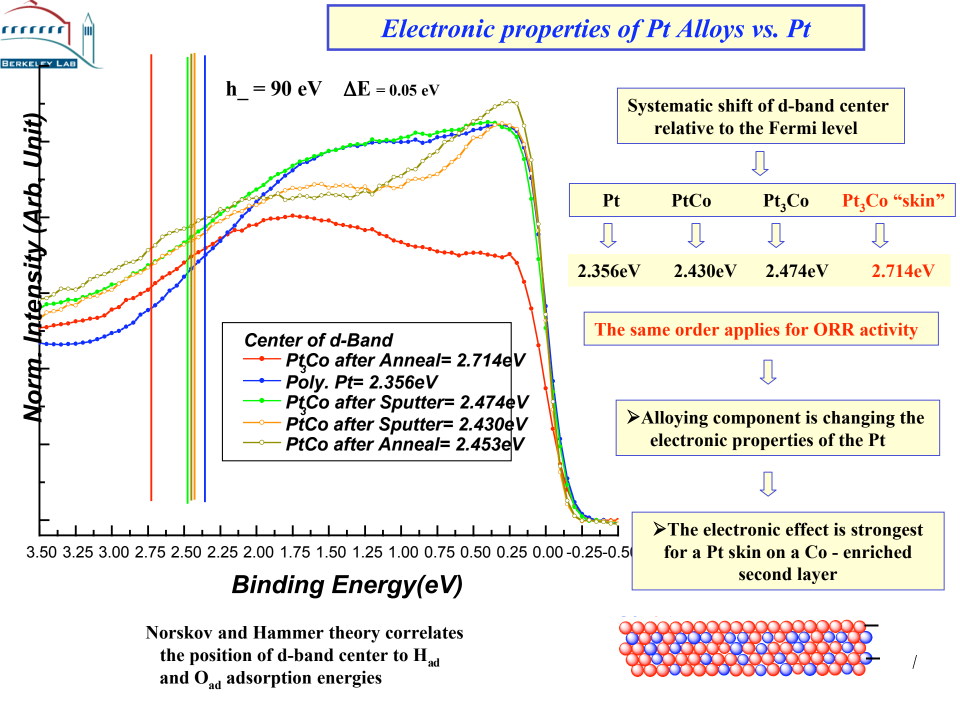
Pt ₃ Ni	Pt ₃ V	Pt ₃ Fe	PtCo	Pt ₃ Co	Pt ₃ Co _{skin}	Pt
~170	XXX	~160	~190	~180	~180	~200



H_{upd} charge is lower but activity for ORR is higher

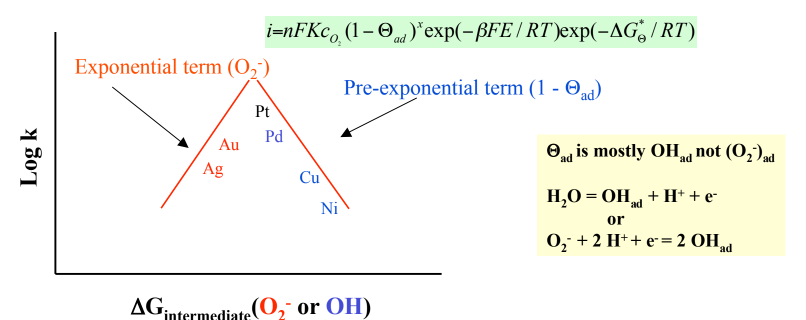


▶ Possible electronic effect of alloying component on Pt skin





The Volcano Relation in ORR Kinetics

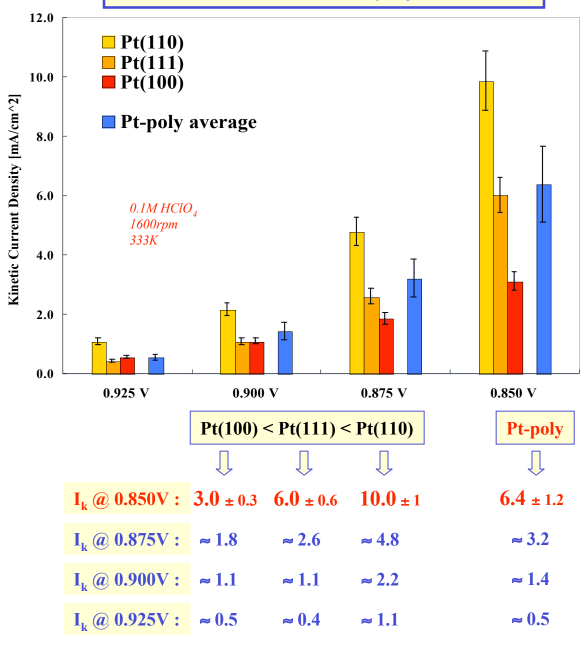


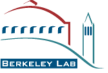
Pt alloys at the Top of the Volcano

- \triangleright Interaction of the electrode with O_2 -requires partially filled d-orbitals with large radial extent Only Group VIII metals satisfy this requirement
- ▶Interaction of the electrode with OH_{ad} must be relatively weak Of the Group VIII metals, Pt has the weakest interaction with OH_{ad} Pt skin on Ni,Co alloy has weaker binding with OH_{ad}

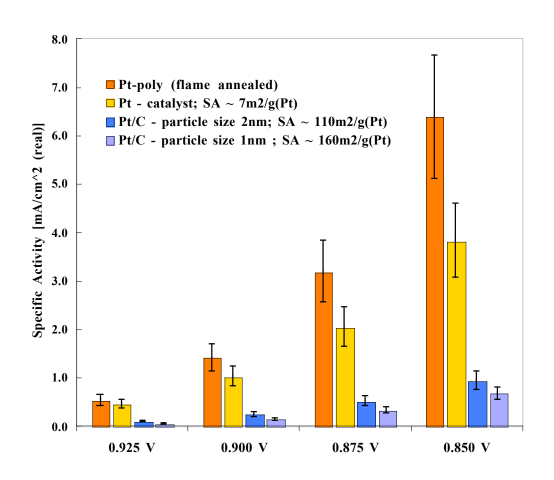


Structure Sensitivity of ORR

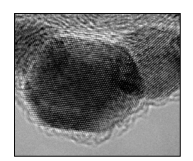


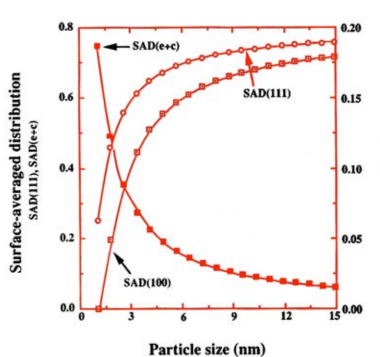


Specific Activity and Particle Size



- Cannot reconcile "loss" of specific activity in Pt nanoparticles with structure sensitivity in single crystals
- There is a maximum in mass activity (mA/mg Pt) at about $60 \text{ m}^2/\text{g}$ (ca. 5 nm particle size)

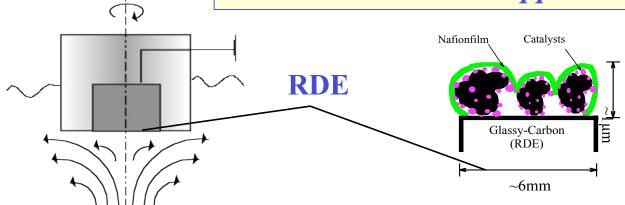






Thin-film RDE method for kinetic measurements with supported catalysts

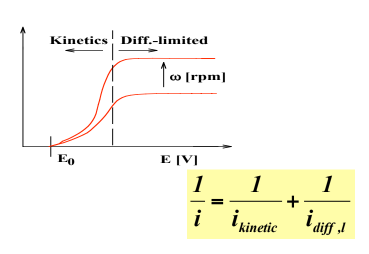




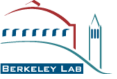
- No agglomerate diffusion
 - > thin catalyst layer, < 1μm
- Negligible mass transport resistance through Nafion film
 - \triangleright 0.1-0.2 µm
- Reproducible loading

$$\gt$$
 $\geq 7\mu g_{metal}/cm_{\perp}$

- **⇒** 100% wetting/utilization
- **→** Fuel cell relevant mass specific current densities

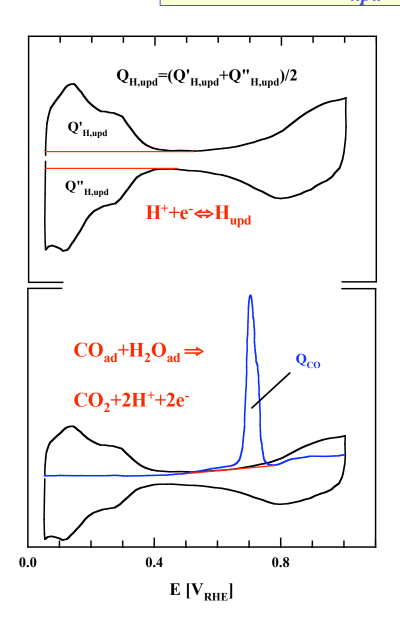


$$i_{diff,l} = 0.62 nFD^{2/3} v^{-1/6} c_{O_2} \omega^{1/2} = B c_{O_2} \omega^{1/2}$$



Roughness factor and real Pt surface from H_{upd} and CO stripping





- **→** Pt loading is expressed over geometric surface area
- \Rightarrow Geometric surface area = 0.283 cm²
- ightharpoonup r_f=1 for Q_{H,upd}=Q_{CO}/2=0.220 mC/cm²

Roughness factor (r_f) and real Pt surface (A_{Pt}) from H_{und} :

$$r_{f} = \frac{Q_{H,upd}}{0.220 mC / cm^{2}} \left[\frac{cm^{2}(real)}{cm^{2}(geo)} \right]$$

$$A_{Pt} = \frac{r_{f}}{Pt_{loading}} \left[\frac{m^{2}}{g_{Pt}} \right]$$

Roughness factor (r_f) and real Pt surface (A_{Pt}) from CO stripping:

$$r_{f} = \frac{Q_{co}/2}{0.220 mC/cm^{2}} \left[\frac{cm^{2}(real)}{cm^{2}(geo)} \right]$$

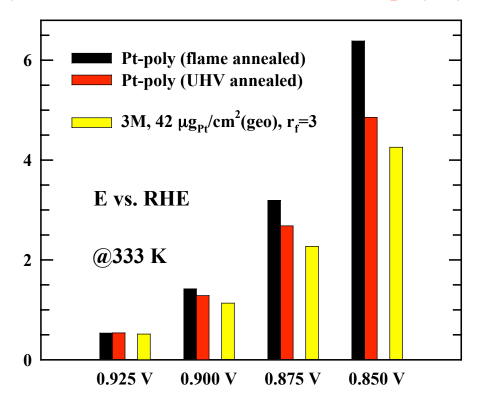
$$A_{Pt} = \frac{r_{f}}{Pt_{loading}} \left[\frac{m^{2}}{g_{Pt}} \right]$$



Summary



- **➡** Thin-film RDE method was optimized for measuring 3M NS Pt catalysts
- \Rightarrow Pt real surface area of 7 10 m²/g_{Pt}
- Activity for ORR close to those obtained on polycrystalline Pt



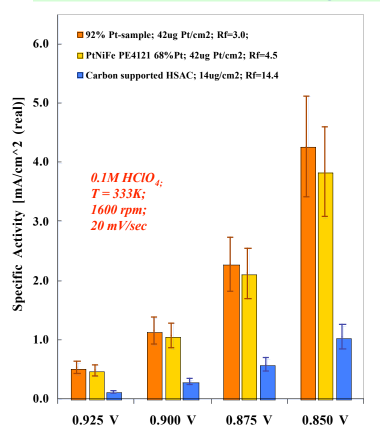
- **□** During ORR no peroxide production in region of mixed kinetic-diffusion control
- → Activation energy of ~22 kJ/mol same as for poly Pt



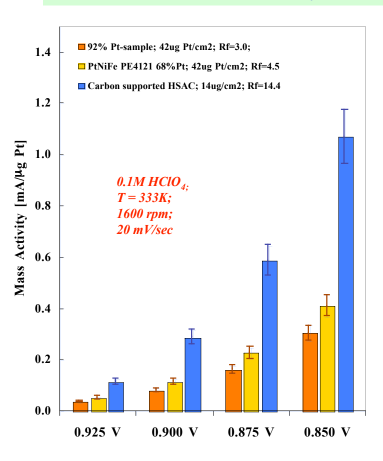
NS Pt Versus Conventional HSAC Supported Pt



(a) Specific Activity [mA/cm²_{geo}]



(b) Mass Activity $[mA/\mu g_{Pt}]$



► Pt alloy NS catalyst has approximately the same specific activity as 92% Pt-sample, but because of the significantly higher roughness factor (Rf=4.5 instead of 3.0) mass activity for alloy is ~35% higher.

► Compared to a carbon-supported High Surface Area Catalyst (Pt-Loading is 14µg/cm²) under same experimental conditions, specific activity is ca. 4 times higher for both 3M-NS catalysts, but mass activity is 3 to 4 times lower.